ALIPHATIC & AROMATIC AMINES

Amines are derivatives of ammonia in which one or more hydrogen atoms are replaced by alkyl group. Amines are classified as primary, secondary or tertiary depending on the number of alkyl groups attached to nitrogen atom. $\mathbb{NH}_{3} \xrightarrow{^{+R}} \to$ $R-NH_2$ $R_{\overline{2}} - NH$ R₃−N s-amine p-amine t-amine (Imino) (Nitrilo) (amino) General formula 1 amine 2 amine 3 amine $C_n H_{2n+1} N H_2$ or $C_n H_{2n+3} N$ $(C_{n}H_{2n+1})_{3}N$ $(C_n H_{2n+1})_2 N H$ Structure : Nitrogen atom is sp³ hybridised. It has one lone pair of electron so the shape of molecule is pyramidal. Isomerism : Amines show chain, position, functional isomerism and metamerism. How many primary amines are possible from molecular formula $C_4H_{11}N$ Ex. (A) 4 (B) 5 (D) 3 (C) 6 Sol. (A) Ex. Which isomerism present in n-propyl amine and isopropyl amine (A) Metamarism (B) Functional group (C) Position (D) Position and chain Sol. (C) How many structural isomers are possible from molecular formula C3HoN Ex. (A) 2 (B) 3 (D) 4 (C) 1 Sol. (B) Write the structures of amines having the molecular formula $C_4H_{11}N$ Ex. Sol. (e) CH_-CH_-NH-CH_CH_ (a) CH-CH-CHNH, CH. $(f) \Box H_3 - N H - \Box H_2 - \Box H_2 - \Box H_3$ (b) CH₃ -attationalist 1 amine $C-NH_2$ d CHάH CH3 | (h) CH3-N-CH2-CH3 CH, 3 amine (d) CH_{-} CH_{-} CH_{-} NH_{2}

General Method of Preparation : q

Ammonolysis of alkyl halides and alcohol : (1)

From Ammonolysis of alkyl halides [Hofmann's ammonolysis] : (a)

When an aqueous solution of ammonia is heated with alkyl halide all the three types of amines and quaternary ammonium salt are formed.

$$R - X \xrightarrow{NH_3} R - NH_2 \xrightarrow{R-X} R_2 - NH \xrightarrow{R-X} R_3N \xrightarrow{R-X} R_4NX$$

(Quaternary ammonium salt)

If ammonia is taken in excess, 1 amine is the main product.

(b) Ammonolysis of alcohols :

When ROH and NH_3 are passed over Al_2O_3 or ThO_2 at 350 C all the three types of amines are formed.

$$R-OH \xrightarrow{NH_3} R-NH_2 \xrightarrow{R-OH} R_2-NH \xrightarrow{R-OH} Al_2O_3 \rightarrow R_3N$$

Note :

Quaternary ammonium hydroxide is not formed due to steric hindrance. (i)

> If excess of ammonia is used, then main product will be primary amine. (ii)

(2)By reduction :

- LiAIH₄ With RCONH₂ : RCONH₂ RCH₂NH₂ (a) or Na/C₂H₅OH
- With RCN : Na/C₂H₅OH (b) RCN + 4HRCH₂NH₂

This reaction is called mendius reaction.

The reduction of alkyl isocynides with sodium and ethanol gives secondary amines.

$$R-NC + 4H$$
 $C_{2H_{5}OH/Na}$ $RNHCH_{3}$

- With Oximes : R-OH-OH+4H $\xrightarrow{\text{LiAlH}_4}$ $\xrightarrow{\text{Na/CaH-OH}}$ RCH,—NH, + H,O (c)
- $\xrightarrow{Sn/HCl}$ $RNH_2 + 2H_2O$ With RNO₂ : $RNO_2 + 6H$ (d)

In lab method we use Sn/HCl while in industrial method we use Fe / HCl.

(3) By hydrolysis of :

R-NC : Alkyl isocyanide undergoes hydrolysis with mineral acid and forms alkyl amine. (a)

 $R-NC + 2H_2O \xrightarrow{HCl} RNH_2 + HCOOH$

RNCO : Alkyl isocyanate undergoes hydrolysis on heating with KOH. (b)

 $R-N=CO + 2 \text{ KOH} \longrightarrow RNH_2 + K_2CO_3$

From Grignard reagent : (4)

Alkyl magnesium iodide reacts with chloramine to yield alkyl amine.

$$R-M_{g}-I + C - NH_{2} \longrightarrow R-NH_{2} + M_{g} \langle I_{C} \rangle$$

(5) Gabriel phthalimide synthesis :

Phthalimide is first treated with KOH to obtain potassium phthalimide which is then treated with alkyl iodide. Then alkyl phthalimide on hydrolysis yields alkylamine. This method is used in the formation of pure aliphatic primary amines.



$$\begin{array}{c} \underset{Acylazide}{\operatorname{Rec}} \xrightarrow{-N_{2}} R \xrightarrow{-N_{2}} R \xrightarrow{-N_{2}} R \xrightarrow{-N_{2}} R \xrightarrow{-N_{2}} R \xrightarrow{-NH_{2}} \\ Alkyl amide \end{array}$$

$$\begin{array}{c} \underset{R \leftarrow C \rightarrow N}{\operatorname{Rec}} R \xrightarrow{-N_{2}} R \xrightarrow{-N_$$

(8) Schmidt reaction :

In presence of conc. $\mathrm{H_2SO_4}$ alkanoic acid reacts with hydrazoic acid (N_3H) to yield alkylamine.

$$R-COOH + N_{3}H \xrightarrow{Conc. H_{2}SO_{4}} R-NH_{2} + N_{2} + CO_{2}$$

Mechanism :



(9) Lossen rearrangement reaction :

In this reaction hydroxamic acid undergoes rearrangement and gives alkyl amine.



 \longrightarrow R-N=C=O $\xrightarrow{H_2O}$ R-NH₂

(10) Reductive amination of aldehyde and ketone :

Physical Properties :

> CH_3NH_2 is gas and $C_2H_5NH_2$ is a volatile liquid. (i)

(ii) Higher amines have fishy smell.

H-Bonding (weaker as compared to H–O –H). (iii)



In 3 amine (due to absence of H-atom) H-bonding is not possible.

(iv) Boiling point : Due to small intermolecular association the b.p. of 1 and 2 amines are lower than those of alcohols of comparable molecular weight. The boiling point of 3 amines which form no H-bonds are near to those of alkanes of comparable molecular weight.

Boiling point α molecular weight

Order of B.P. :	1	amine	>	2	amine	>	3	amine
so order of volatility :	3	amine	>	2	amine	>	1	amine

(v) **Solubility** : Low molecular weight amines (< six carbon) are very soluble in water. The water solubility of amines decreases with increasing size of alkyl group.



□ Chemical Properties :

- (i) Basic character of amines is due to the presence of lone pair of electrons on the N atom.
- (ii) Basic strength depends on electron donating tendency.

Basicity order in aqeous solution and in liquid phase. $Et_2 NH > Et_3 N > Et NH_2$. Due to steric hindrance in 3 amine, it is less basic, than 2 amine.

Steric hindrance of three $-C_2H_5$ group protect the lone pair of nitrogen from the attack of H^{\oplus} .

But in gaseous phase basic order is $R_2NH > RNH_2 > R_3N > NH_3$

Some other basic order of different amine if alkyl group would be change

- Alkyl groups (R–) Relative base strength
- (i) $CH_3 R_2NH > RNH_2 > R_3N > NH_3$
- (ii) $C_2H_5 R_2NH > RNH_2 > NH_3 > R_3N$
- (iii) $(CH_3)_2CH RNH_2 > NH_3 > R_2NH > R_3N$
- (iv) $(CH_3)_3C$ $NH_3 > RNH_2 > R_2NH > R_3N$

□ Special point :

- (I) Tertiary amine is less basic then secondary due to following reasons :
- (i) **Steric hindrance**: In tertiary amines (R_3N) , three alkyl groups attached to N are bulkier and as such exert steric hindrance.
- (ii) Decrease in hydration :

In tertiary amine

 $R_{s}^{\oplus} H - H - K$

In secondary amine

Protonated t-amine can form H-bonding with water molecule only at one point [less stable] Protonated s-amine can form H-bonding with water molecules at two points (more stable)

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3 amine are less stable as compare to 2 amine due to low hydration so less basic.

(II) The basic strength of aniline is less than aliphatic amines as the lone pair of electron present on N- atom interact with the delocalized π - orbital of benzene ring. Hence it is less available for protonation on N-atom.

> Aniline

 $\xrightarrow{\Delta} R-Cl + NH_3$

>

Pyrrole

The basic order nature for aniline, pyridine and pyrrole \longrightarrow Pyridine

(2) Reactions showing basic nature :

(a) It reacts with acids to form salts.

 $RNH_2 + HCl \longrightarrow [RNH_3] \overset{\oplus}{\cap}$

Alkyl ammonium chloride

(Acidic salt)

$$2RNH_2 \xrightarrow{H_2SO_4} (RNH_3)_2 SO_4^{-2} Alkyl ammonium sulphate$$

(b) Amines reacts with auric acid and platinic chlorides in presence of HCl to form double salts. These double salts decompose on ignition to pure metal. Therefore the formation and decomposition of the double salts is used for determining the molecular weight of amines.

$$2R - NH_{2} + PtQ_{4} + 2HQ \longrightarrow R^{\oplus} H_{3}PtQ_{6}^{-2}$$
(chloro platinic acid) Alkyl ammonium chloroplatinate
$$RNH_{2} + ALQ_{3} + HQ \longrightarrow R^{\oplus} H_{3}AUQ_{4}$$
[Chloroauric acid] Alkyl ammonium chloroaurate
$$(R^{\oplus} H_{3})_{2} PtCl_{6}^{-2} \longrightarrow Pt$$

(c) Reaction with H_2O : It forms alkyl ammonium hydroxide with water ammonium hydroxides are used for precipitation of IInd and IIIrd group cations in qualitative analysis

$$RNH_{2} + H_{2}O \longrightarrow (RNH_{3})OH Base$$

$$FeCl_{3} + 3[RNH_{3}]OH \longrightarrow Fe(OH)_{3} + 3[RNH_{3}]Cl$$

$$Brown ppt$$

$$AICl_{3} + 3[RNH_{3}]OH \longrightarrow AI(OH)_{3} + 3[RNH_{3}]Cl$$

$$White ppt.$$

$$CrCl_{3} + 3[RNH_{3}]OH \longrightarrow Cr(OH)_{3} + 3[RNH_{3}]Cl$$

$$Green ppt.$$

(3) Reaction with alkyl halides :

Alkyl amine reacts with alkyl halides and form sec., ter. amines and quaternary ammonium salt.

 $RNH_2 + R-X \xrightarrow{-HX} R_2NH \xrightarrow{-RX} R_3N \xrightarrow{RX} R_4NX$

Special Point :

Separation of 1, 2 and 3 amines :

1, 2, 3 amine + $R_a NX$ $\xrightarrow{distillation}$ Mixture of 1, 2, 3 amine

 R_4NX does not undergo distillation.

Mixture of 1, 2, 3 amine can be separated by following methods.

- (i) Fractional distillation : The mixture of amines may be separated by fractional distillation because their boiling points are quite different. It is used in industry.
- (ii) Hinsberg method : In this method mixture of amines is seperated by using benzene sulphonyl chloride (Hinsberg's reagent).

 $C_6H_5SO_2Cl + 1$ amine \longrightarrow Product \xrightarrow{KOH} dissolve

 $C_{c}H_{r}SO_{2}Cl + 2$ amine \longrightarrow Product \xrightarrow{KOH} insoluble

3 amine does not react with benzene sulphonyl chloride.

- (iii) Hofmann method : In this method mixture of amines is separated by using ethyl oxalate.
 - 1 amine + ethyl oxalate \longrightarrow solid product
 - 2 amine + ethyl oxalate \longrightarrow liquid product
 - 3 amine + ethyl oxalate \longrightarrow No reaction
- (4) Acetylation : Acetylation takes place when alkyl amine combines with acetyl chloride or acetic anhydride.

$$RNH_{2} + ClCOCH_{3} \longrightarrow RNHCOCH_{3} + HCl$$

$$(N -alkyl acetamide)$$

$$RNH_{2} + (CH_{3}CO)_{2}O \longrightarrow RNHCOCH_{3} + CH_{3}COOH$$

$$(N -alkyl acetamide)$$

(5) Benzoylation (Schotten baumann reaction) :



(6) Acidic nature : Amines are very weak acids only 1 and 2 amines show acidic nature.

$$H_2 + Na \longrightarrow RNHNa + \frac{1}{2}H_2$$

N- alkyl sodamide

(7) Reaction with Tilden reagent :

When alkylamine reacts with nitrosyl chloride (Tilden reagent) alkyl chloride is formed. This reaction is important in interconversion.

 $R-NH_2 + NOCl \longrightarrow RCl + N_2 + H_2O$

(8) Reaction with phosgene :

 $R-NH_2 + COCl_2 \longrightarrow$

$$R = N = C = O + 2HC$$

Alkyl isocyanate

(9) **Reaction with halogen :** The hydrogen atoms of the amino group are replaced by halogen atoms in presence of alkali solution.

$$R-NH_2 + Br_2 \xrightarrow{NaOH} R-NH-Br + HBr$$

N– bromo alkylamine

$$R-NH-Br + Br_2 \xrightarrow{NaOH} R-NBr_2 + HBr$$

N, N-dibromo alkyl amine

(10) Reaction with aldehydes : Alkylamine reacts additively with aldehydes to form α - hydroxyl amines which are changed to schiff bases with elimination of water molecule.

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$$R-NH_{2} + O=C-R \longrightarrow R-N-C-R \longrightarrow R-H_{2}O \xrightarrow{-H_{2}O} RCH=NR (Schiff's base)$$

(11) Mannich reaction :

$$R-CH_2NH_2 + CH_2O + HCH_2-C-C_6H_5 \xrightarrow{-H_2O} R-CH_2NHCH_2-C+C_6H_5$$

Acetophenone Mannich base

(12) Oxidation :

 KMnO₄/H⁺: Alkylamine on oxidation with acidified potassium permaganate forms aldimine which on hydrolysis gives aldehyde and ammonia.

$$R-CH_2NH_2 \xrightarrow{[0]} R - CH = NH \xrightarrow{H_2O}_{H^+} RCHO + NH_3$$
(Alclimine)

$$R_{2}CHNH_{2} \xrightarrow{[0]} R_{2}C = NH \xrightarrow{H_{2}O} R_{2}C = O + NH_{3}$$
(Ketimine) (Ketone)

$$R_3CNH_2$$
 R_3CNO_2 (Nitroalkane)

• With H_2SO_5 (Caro's acid) Or H_2O_2 /Fe⁺² (Fenton reagent) :

When alkyl amine's heated with chloroform and alc. KOH alkyl isocyanide is formed which has very bad smell.

This test is also given by aniline . This is a test for p- amines.

$$R-NH_2 + CHCl_3 + 3 \text{ KOH} \longrightarrow R - N \equiv C + 3KCl + 3H_2O$$

Nucleophile RNH_2 attacks electrophilic intermediate [: CCl_2] dichlorocarbene.

Mechanism :



When alkyl amine is heated with carbon disulphide and mercuric chloride alkyl isothiocyanate is formed which has smell like mustard oil.



- (ii) Amines can have chiral N-atom but cannot be resolved into enantiomeric forms because of rapid inversion of one enantiomeric form into the other.
- (iii) The Schiff's bases formed by reaction of 1 -amines and aldehyde/ketones are also called anils.
- y) The mixture of 1, 2, 3 amines can be distinguished by Hoffmann's test or Hinsberg's reagent or carbylamine test or nitrous acid test.
- (v) In Hoffmann test $CS_2 + HgCl_2$ are used and in Hinsberg test benzene sulphonyl chloride ($C_6H_5SO_2Cl$) is used.

ANILINE $(C_6H_5NH_2)$

Aniline is also called aminobenzene or phenyl amine. Aniline was first prepared by **Unverdon** 1826 by the distillation of indigo which is called **anil** in spanish and hence the name aniline. In aniline $-NH_2$ group is directly attached to benzene ring.

- General Methods of Preparation :
- (1) Lab method : Aniline is prepared in the lab by reduction of $C_6H_5NO_2$ with Sn + HCl.

$$C_6H_5NO_2 \xrightarrow{(i)Sn+HCl} C_6H_5NH_2 + H_2O$$

(2) Industrial method : Aniline is obtained by reduction of nitrobenzene in presence of Fe/HCl.

$$C_6H_5 - NO_2 \xrightarrow{(i) Fe+HCl} C_6H_5 - NH_2 + H_2O$$

(3) From Phenol : Aniline is obtained when phenol is treated with ammonia in presence of ZnCl₂ at 300 C.

$$C_6H_5OH + NH_3 \xrightarrow{ZnCl_2} C_6H_5NH_2 + H_2O$$

(4) From benzamide (Hofmann's reaction): Aniline is formed when benzamide is treated with Br₂ and KOH.

$$C_6H_5CONH_2 + Br_2 + 4KOH \longrightarrow C_6H_5NH_2 + K_2CO_3 + 2KBr + 2H_2O$$

(5) From benzoic acid (Schmidt reaction) : Benzoic acid is dissolved in conc. H_2SO_4 and hydrazoic acid is dissolved in chloroform. When both solutions are mixed aniline obtained.

$$C_6H_5COOH + N_3H$$

(Hydrazoic acid) $\xrightarrow{Conc.} C_6H_5NH_2 + N_2\uparrow + CO_2\uparrow$

(6) From chloro benzene : Aniline can be manufactured by the action of ammonia on chloro benzene in presence of cuprous oxide (Cu_2O).

$$2C_6H_5Cl + Cu_2O + 2NH_3 \longrightarrow 2 C_6H_5NH_2 + 2CuCl + H_2O$$

(7) From Grignard reagent :

$$C_6H_5MgBr + CINH_2 \longrightarrow C_6H_5NH_2 + Mg A_0$$

(8) From Benzene:

$$C_6H_6 + NH_2OH \longrightarrow C_6H_5NH_2 + H_2OH$$

(9) From phenyl isocyanide :

 $C_{6}H_{5}N \cong C + 2H_{2}O \longrightarrow C_{6}H_{5}NH_{2} + HCOOH$

(10) From phenyl isocyanate:-

 $C_6H_5N = C = O + 2KOH \longrightarrow C_6H_5NH_2 + K_2CO_3$

- Physical Properties :
 - Fresh, aniline is a colourless oily liquid. On standing the colour becomes dark brown due to action of air and light.
 - It's B.P. is 183 C.
 - (iii) It is heavier than water.
 - (iv) It has characteristic unpleasent odour. It is toxic in nature.

□ Similarities and Differences between Aromatic and Aliphatic amines :

(A) Similarities :

- (i) Both are basic, although aliphatic amines are more basic than the aromatic amines.
- (ii) Both form salts with acids, however salts of aromatic amines are easily hydrolysed.
- (iii) Both undergoes alkylation and acylation.
- (iv) Both react with Grignard reagents forming hydrocarbons.
- (v) Both forms schiff's bases.

(B) Differences :

- (i) Aniline is insoluble in water while aliphatic amines are soluble in water (due to H-bonds)
- (ii) Aniline gives diazonium salt with HNO₂ while aliphatic amines gives alcohol and nitrogen (except CH₃NH₂)

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- (iii) Aniline undergoes coupling and electrophilic substitution reactions in benzene ring while aliphatic is not.
- (iv) Aniline has characteristic aromatic smell while aliphatic amines have smell like ammonia
- (v) Aniline gives aniline black dye with acidic $K_2Cr_2O_7$ while aliphatic does not form dye.
- (vi) Aniline gives violet colour with NaOCl while aliphatic amines does not give.

□ Chemical Properties :

- (i) Aniline is a primary amine it shows properties of both of benzene nucleus and $-NH_2$ group.
- (ii) Aniline has weak basic nature as compared to aliphatic amine.
- (ii) Weaker basic nature of aniline as compared to aliphatic amines can be explained on the basis of resonance. In aniline the non-bonding electron pair of N is delocalised in to benzene ring by resonance. Thus electron density is less on N atom due to which aniline is less basic than aliphatic amines

Order of basic strength : $RNH_2 > NH_3 > C_6H_5NH_2$

(iv) Electron withdrawing group decreases the basic strength where as electron donating groups increases the basic strength

$$p - C_{e}H_{A}$$
 NO_{2} $C_{6}H_{5}NH_{2}$ $p - C_{e}H_{4}$ NH_{2} NH_{2}

- (v) Aqueous solution of aniline is neutral to litmus
- (A) Reactions due to -NH₂ group :
- (1) Basic nature Aniline is weak base but it forms salt with strong acids. It a accepts a proton.

Chloro platinic method : This is used to determination of molecular weight of organic compounds.

Ex. Conjugate base of $(CH_3)_2 \mathbb{NH}_2^{\oplus}$ is

(A)
$$(CH_3)_3 N$$
 (B) $(CH_3)_2 NH$ (C) $(CH_3)_2 N^{\oplus}$ (D) $(CH_3)_2 N^{\oplus}$

Sol. (B)

(2) Alkylation :

Aniline reacts with alkyl halides forming secondary, tertiary and quaternary ammonium salts depending on the concentration of alkyl halides.

(3) Acylation :

Aniline reacts with acid chlorides or anhydrides to form corresponding amides called anilides. The reaction of $C_6H_5NH_2$ with benzoyl chloride is called "Schotten Baumann reaction".

$$C_{6}H_{5}NH_{2} + Q - C - QH_{3} \longrightarrow C_{6}H_{5} - NH - C - QH_{3} \quad (Acetanilide)$$

$$C_{6}H_{5}NH_{2} + (QH_{3} - C -)_{2}O \longrightarrow C_{6}H_{5} - NH - C - QH_{3} + CH_{3}COOH$$

$$C_{6}H_{5}NH_{2} + Q - C - C_{6}H_{5} \longrightarrow C_{6}H_{5} - NH - C - C_{6}H_{5} \quad (Benzanilide)$$

$$C_{6}H_{5}NH_{2} + Q - C - C_{6}H_{5} \longrightarrow C_{6}H_{5} - NH - C - C_{6}H_{5} \quad (Benzanilide)$$

(4) **Carbylamine reaction**: When aniline is heated with CHCl₃ and KOH it gives isocyanide having unpleasent smell which can be easily detected.

$$C_6H_5NH_2 + CHCl_3 + 3KOH \longrightarrow C_6H_5NC + 3KCl + 3H_2O$$

Phenyl isocyanide

• Note : (i) Intermediate species is dichloro carbene [: CCl₂].

This is a test of aniline and other primary amines and is known as isocyanide test.

(5) Hoffmann's mustard oil reaction :

(ii)

When aniline is heated with alc. CS_2 and excess of $HgCl_2$ phenyl isothiocyanate having a characteristic smell of mustard oil is formed.

$$C_6H_5NH_2 + S = C = S \xrightarrow{H_gCl_2} C_6H_5N = C = S$$

Phenyl isothiocyanate

If the above reaction is carried out in presence of solid KOH diphenyl thiourea is formed (Used as accelerator for vulcanisation of rubber).

$$\begin{array}{ccc} C_{6}H_{5}NH_{2} \\ C_{6}H_{5}NH_{2} \end{array} + S=C=S \longrightarrow \begin{array}{ccc} C_{6}H_{5}NH \\ C_{6}H_{5}NH_{2} \end{array} C=S + H_{2}S \end{array}$$

When diphenyl thiourea is treated with HCl it gives phenyl isothiocyanate.

$$\begin{array}{c} C_{6}H_{5}NH\\ C=S + HO \longrightarrow C_{6}H_{5}-N=C=S + C_{6}H_{5}NH_{3}CI \end{array}$$

Reaction with aldehydes : Aniline condenses with aldehydes to form schiff's base. (6)

$$C_6H_5NH_2 + H - C - C_6H_5 \longrightarrow O$$

 $C_{H}N = C_{H}C_{H}H_{2} + H_{2}O$ Benzylidene aniline (schiff's base)

(7)Reaction with Heinsberg's reagent :

$$C_6H_5 - SO_2Cl + HNHC_6H_5 \xrightarrow{-HCl} C_6H_5SO_2NHC_6H_5$$

(N- Phenylbenzene sulphonamide)

(8)Acidic nature :

$$C_6H_5NH_2 + Na \longrightarrow C_6H_5NHNa + 1/2H_2$$

N-Phenyl sodamide

(9) Diazotisation :

Diazotisation is a reaction in which ice cooled solution of aniline in a inorganic acid with sodium nitrite solution leading to the formation of diazonium salt.

$$C_{6}H_{5}NH_{2} + NaNO_{2} + HCI \xrightarrow{0^{\circ}-5^{\circ}C} -H_{2}O \rightarrow C_{6}H_{5}N_{2}CI$$

$$C_{6}H_{5}NH_{2} + HCI \xrightarrow{0^{\circ}-5^{\circ}C} -H_{2}O \rightarrow C_{6}H_{5}N_{2}CI$$

$$NaNO_{2} + HCI \xrightarrow{0^{\circ}-5^{\circ}C} -H_{2}O \rightarrow C_{6}H_{5}N_{2}CI + 2H_{2}O$$
Benzene diazonium chloride

Benzene diazonium chloride is a useful synthetic reagent. It is used in the preparation of many organic compounds

Note :

Coupling between arenediazonium cations and amines take place most rapidly in slightly acidic solution (pH 5 to 7). Under these conditions the concentration of the arenediazonium cation is at a maximum ; at the same time an excessive amount of the amine has not been converted to an unreactive aminium salt.

$$C_{6}H_{5}N_{2}Cl \xrightarrow{C_{6}H_{5}OH} C_{6}H_{5}N=N-C_{6}H_{4}OH \text{ [p-hydroxy azobenzene (orange dye)]}$$

$$C_{6}H_{5}N_{2}Cl \xrightarrow{C_{6}H_{5}NH_{2}} C_{6}H_{5}N=N-C_{6}H_{4}NH_{2} \text{ [p-amino azobenzene (aniline yellow)]}$$

$$C_{6}H_{5}N_{2}Cl \xrightarrow{\beta-Naphthol} OH \text{ (Red colour)}$$

(10) Oxidation : Aniline forms a number of products depending upon the nature of oxidising agent:-

 $-NH_2 \xrightarrow{[0]} -NHOH \xrightarrow{[0]} -NO \xrightarrow{[0]} -NO_2$

S.N.	Oxidant		Product
1.	Acidic KMnO ₄		Aniline black (a dye)
2.	Alkaline KMnO ₄		Azobenzene $C_6H_5N = NC_6H_5$
3.	Neutral KMnC	04	Azobenzene + Nitro benzene
4.	Caro's acid (H	₂ SO ₅)	Nitrobenzene + Nitroso benzene
5.	CF ₃ COOOH		Nitrobenzene
6.	$K_2 Cr_2 O_7 + co$	nc. H_2SO_4	p-Benzo quinone
7.	NaOCl		p- Amino Phenol (Violet colour)
3.	HNO ₃		Decomposes
€.	Aniline $\xrightarrow{Atmospheric air} \& light$		Dark red colour
	C_2H_5OH	7	
	$H_3PO_2 \rightarrow$		
		$C_{6}H_{6}$ Benzene	S.
	$\xrightarrow{\text{Na/NaOH}} 2\text{H}$		69
	$\xrightarrow{\operatorname{Cu}_2\operatorname{Cu}_2/\operatorname{HC}}$	$C_6H_2O + N_2 + HO$	
	$\xrightarrow{\text{Cu}_2\text{Br}_2/\text{HBr}} \rightarrow$	$C_{e}H_{s}Br + N_{2} + HC$	Sandmeyer reactions
	$\xrightarrow{\operatorname{Cu}_2(\operatorname{CN})_2/\operatorname{HCN}}$	C₀H₄CN + №₂ + HCI	
GHN₂($\square \xrightarrow{KI (aq.)} \rightarrow$	C₀HၞI + № + KCI	
	NaBF₄∕∆ Sodium tetrafluoro borate	$C_6H_3F + BF_3 + NaCl$	Balz-Schiemann reaction
	H ₂ O	CHOH	
		C,H,O	Gattermann reactions
	$\xrightarrow{+4H}$ SnCl ₂ +HCl	C6H3NHNH2(Phenyl hy N2-C6H3	xtrazine)
	β−Naphthol 10% NaOH	OO (Red o	xolour)
N	SnCl ₂ /NaOH	C ₆ H ₆	
	$C_6H_0OH \rightarrow$	$C_{c}H_{t}N=N-C_{c}H_{4}OH$	
•	$C_6H_5NH_2$	p-Hydroxy azobenzene $C_{6}H_{5}N=N-C_{6}H_{4}NH_{2}$	(Orange dye) Coupling reactions



- Note :
 - (i) In aniline 2, 4, 6 or ortho and para positions are electron rich so electrophile attacks here. In aniline 3, 5, or meta position is electron deficient so nucleophile attacks here.
 - (ii) The benzene ring of aniline undergoes halogenation, sulphonation and nitration.
 - (iii) The NH_2 group is o-, p-directing.
- (1) Halogenation : Chlorine and bromine react with aniline and form trichloro and tribromo aniline respectively



Note :

However, monobromo or chloro derivative of aniline can be prepared if $-NH_2$ group is first protected by acetyl group. Here the reactivity decreases due to -I effect of acetyl group.



(2) Nitration :

(a) Direct nitration : The direct nitration of aniline by conc. HNO_3 and conc. H_2SO_4 give meta-nitroaniline. Due to positively charged N, m-position becomes electron rich as compared to o, p-position.



(b) Indirect nitration : In indirect nitration amino group is protected by acetylation to give acetanilide, which on nitration and subsequent hydrolysis give o- and p- nitro-aniline.



- Ex. Azo dye test is given by
 - (A) All amines

- (B) Only secondary amine
- (C) Only primary aliphatic amine (D) Only primary aromatic amine

Sol. (D)

(3) Sulphonation : Aniline reacts with fuming H_2SO_4 to give sulphanilic acid.(p-Amino-benzene sulphonic acid)



- Note : (i) This process is called baking.
 - (ii) Sulphanilic acid is an important intermediate in the manufacturing of dyes and drugs.
 - (iii) The compounds in which both proton donating & proton accepting groups present are called ampholite (dipolar ion).



(4) Catalytic hydrogenation :

Aniline undergoes hydrogenation in presence of Ni at high temp. to form amino cyclohexane.





o– Amino phenyl mercuric acetate

□ Tests of aniline :

• **Carbylamine test** : Aniline gives carbylamine test or Isocyanide test.

$$C_6H_5NH_2 + CHCl_3 + KOH \longrightarrow C_6H_5NC$$

(Bad smelling)

- Dye test : Aniline is first diazotised. On adding alkaline soln. of β-naphthol to the diazotised product a scarlet red dye is formed.
- On heating with bromine water, a ppt. is formed.

Uses of Aniline : Aniline is used in

- (i) The manufacture of dyes and dye intermediates
- (ii) The manufacture of accelerators and antioxidants in rubber industry.
- (iii) The manufacture of acetanilide, sulphuric acid and indigo
- (iv) The Manufacture of sulpha drugs.

SOLVED EXAMPLES





- **9**. A compound X with seven carbon atoms on treatment with Br₂ and KOH gives Y. Y gives carbylamine test and upon diazotisation and coupling gives azodye. X is :
 - (A) C₆H₅CONH₂

(B) $CH_3 - O - C_6H_4NH_2$

- **Sol.** Since Y gives coupling reaction after diazotization, it suggest that Y can be aniline or benzene ring substituted aniline. Since Y has been obtained from Hofmann bromamide it means it has $-\text{CONH}_2$ group with benzene ring. Hecne, it is $C_6H_5\text{CONH}_2$. Hence (A) is the correct answer.
- **10.** Methyl ethyl propyl amine forms non-superimposable mirror images but it does not show optical activity because :

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(A) of rapid flipping
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(B) amines are basic in nautre

(C) nitrogen has a lone pair of electrons

- (D) of absence of asymmetric nitrogen
- **Sol.** The interconversion of d and l-forms are so fast that it is not possible to isolate these. Hence, (A) is the correct answer.

Rapid